LUNAR: Cellular Automata for Drifting Data Streams

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Abstract

With the advent of huges volumes of data produced in the form of fast streams, realtime machine learning has become a challenge of relevance emerging in a plethora of real-world applications. Processing such fast streams often demands high memory and processing resources. In addition, they can be affected by non-stationary phenomena (concept drift), by which learning methods have to detect changes in the distribution of streaming data, and adapt to these evolving conditions. A lack of efficient and scalable solutions is particularly noted in real-time scenarios where computing resources are severely constrained, as it occurs in networks of small, numerous, interconnected processing units (such as the so-called Smart Dust, Utility Fog, or Swarm Robotics paradigms). In this work we propose LUNAR, a *streamified* version of cellular automata devised to successfully meet the aforementioned requirements. It is able to act as a real incremental learner while adapting to drifting conditions. Extensive simulations with synthetic and real data will provide evidence of its competitive behavior in terms of classification performance when compared to long-established and successful online learning methods.

Keywords: Cellular automata, real-time analytics, data streams, concept drift

1. Introduction

Real-Time Analytics (RTA), also referred to as stream learning, acquired special relevance years ago with the advent of the Big Data era (Laney, 2001; Bifet, Gavaldà, Holmes & Pfahringer, 2018), becoming one of its most widely acknowledged challenges. Data streams are the basis of the real-time analytics, composed by sequences of items, each having a timestamp and thus a temporal order, and arriving one by one. Due to the incoming sheer volume of data, real-time algorithms cannot explicitly access all

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historical data because the storage capacity needed for this purpose becomes unmanageable. Indeed, data streams are fast and large (potentially, infinite), so information must be extracted from them in real-time. Under these circumstances, the consumption of limited resources (e.g. time and memory) often implies sacrificing performance for efficiency of the learning technique in use. Moreover, data streams are often produced by non-stationary phenomena, which imprint changes on the distribution of the data, leading to the emergence of the so-called *concept drift*. Such a drift causes that predictive models trained over data flows become eventually obsolete, and do not adapt suitably to new distributions. Therefore, these predictive models need to be adapted to these changes as fast as possible while maintaining good performance scores (Gama, Žliobaitė, Bifet, Pechenizkiy & Bouchachia, 2014).

For all these reasons, the research community has devoted intense efforts towards the development of Online Learning Methods (OLMs) capable of efficiently undertaking predictive tasks over data streams under minimum time and memory requirements (Widmer & Kubat, 1996; Ditzler, Roveri, Alippi & Polikar, 2015; Webb, Hyde, Cao, Nguyen & Petitjean, 2016; Lu, Liu, Dong, Gu, Gama & Zhang, 2018; Losing, Hammer & Wersing, 2018; Lobo, Del Ser, Bifet & Kasabov, 2020). The need for overcoming these setbacks stems from many real applications, such as sensor data, telecommunications, social media, marketing, health care, epidemics, disasters, computer security, electricity demand prediction, among many others (Žliobaitė, Pechenizkiy & Gama, 2016). The Internet of Things (IoT) paradigm deserves special attention at this point (Manyika, Chui, Bisson, Woetzel, Dobbs, Bughin & Aharon, 2015; De Francisci Morales, Bifet, Khan, Gama & Fan, 2016), where a huge quantity of data is continuously generated in real-time by sensors and actuators connected by networks to computing systems. Many of these OLMs are based on traditional learning methods (i.e. Naive Bayes, Support Vector Machines, Decision Trees, Gaussian methods, or Neural Networks (Bifet, Gavaldà, Holmes & Pfahringer, 2018)), which have been streamified to make them work incrementally and fast. By contrast, other models are already suitable for mining data streams by design (Cervantes, Gagné, Isasi & Parizeau, 2018). Unfortunately, most existing RtML models show a high complexity and dependence on the value of their parameters, thereby requiring a costly tuning process (Losing, Hammer & Wersing, 2018). In addition, some of them are neither tractable nor interpretable, which are features lately targeted with particularly interest under the eXplainable Artificial Intelligence (XAI) paradigm (Barredo Arrieta, Díaz-Rodrguez, Del Ser, Bennetot, Tabik, Barbado, García, Gil-López, Molina, Benjamins, Chatila & Herrera, 2019). Nowadays, learning algorithms featuring these characteristics are still under active search, calling for new approaches that blow a fresh breeze of novelty over the field (Gama, Žliobaitė, Bifet, Pechenizkiy & Bouchachia, 2014; Khamassi, Saved-Mouchaweh, Hammami & Ghédira, 2018).

Cellular Automata (CA) become fashionable with the Conway's Game of Life in 1970, but scientifically relevant after the Stephen Wolfram's study in 2002 (Wolfram, 2002). Despite they are not widely used in data mining tasks, the Fawcett's work (Fawcett, 2008) showed how they can turn into a simple and low-bias data mining

method, robust to noise, and with competitive classification performances in many cases. Until now, their appearance on the RTA has been timid, without providing evidences of their capacity for incremental learning and drift adaptation.

This work enters this research avenue by proposing the use of Cellular Automata for RTA. Our approach, hereafter coined as *celluLar aUtomata for driftiNg dAta stReams* (LUNAR), capitalizes on the acknowledged capacity of CA to model complex systems from simple structures. We show the intersection of CA and RTA in the presence of concept drift, showing that CA are promising incremental learners capable of adapting to evolving environments. Specifically, we provide a method to transform a traditional CA into its *streamified* version (sCA) so as to learn incrementally from data streams. We show that LUNAR performs competitively with respect to other online learners on several real-world datasets. More precisely, we will provide informed answers to the following research questions:

- RQ1: Does sCA act as a real incremental learner?
- RQ2: Can sCA efficiently adapt to evolving conditions?
- RQ3: Does our LUNAR algorithm perform competitively with respect to other consolidated RTA approaches reported in the literature?

The rest of the manuscript is organized as follows: first, Section 2 provides a general introduction to CA, placing an emphasis on their historical role in the pattern recognition field, and concretely their relevance for stream learning. Next, Section 3 delves into the methods and the LUNAR approach proposed in this work. Section 4 introduces the experimental setup, whereas Section 5 presents and discusses the obtained results from such experiments. Finally, Section 6 draws conclusions and future research lines related to this work.

2. Related Work

Before going into the technical details of the proposed approach, we herein provide a historical overview of CA (Subsection 2.1), along with a perspective on how these models have been progressively adopted for pattern recognition (Subsection 2.3) and, more lately, stream learning (Subsection 2.4).

2.1. The Limelight Shone Down on Cellular Automata

The journey of CA was initiated by John von Neumann (Neumann, Burks et al., 1966) and Ulam for the modeling of biological self-reproduction. They became really fashionable due to the popularity of Conway's Game of Life introduced by Gardner (Games, 1970) in the field of artificial life (Langton, 1986). Arguably, the most scientifically significant and elaborated work on the study of CA arrived in 2002 with the thoughtful studies of Stephen Wolfram (Wolfram, 2002). In recent years, the notion of complex systems proved to be a very useful concept to define, describe, and study

various natural phenomena observed in a vast number of scientific disciplines. Despite their simplicity, they are able to describe and reproduce many complex phenomena (Wolfram, 1984) that are closely related to processes such as self-organization and emergence, often observed within several scientific disciplines: biology, chemistry and physics, image processing and generation, cryptography, new computing hardware and algorithms designs (i.e., automata networks (Goles & Martínez, 2013) or deconvolution algorithms (Zenil, Kiani, Zea & Tegnér, 2019)), among many others (Ganguly, Sikdar, Deutsch, Canright & Chaudhuri, 2003; Bhattacharjee, Naskar, Roy & Das, 2016). CA have also been satisfactorily implemented for heuristics (Nebro, Durillo, Luna, Dorronsoro & Alba, 2009) and job scheduling (Xhafa, Alba, Dorronsoro, Duran & Abraham, 2008). Besides, the capability to perform universal computation (Cook, 2004) has been one of the most celebrated features of CA that has garnered the attention of the research community: an arbitrary Turing machine can be simulated by a cellular automaton, so universal computation is possible (Wolfram, 2002).

Considering their parallel nature (which allows special-purposed hardware to be implemented), CA are also called to be a breakthrough in paradigms such as Smart Dust (Warneke, Last, Liebowitz & Pister, 2001; Ilyas & Mahgoub, 2018), Utility Fog (Hall, 1996; Dastjerdi & Buyya, 2016), Microelectromechanical Systems (MEMS or "motes") (Judy, 2001), or Swarm Intelligence and Robotics (Ramos & Abraham, 2003; Del Ser, Osaba, Molina, Yang, Salcedo-Sanz, Camacho, Das, Suganthan, Coello & Herrera, 2019), due to their capability to be computationally complete. Microscopic sensors are set to revolutionize a range of sectors, such as space missions (Niccolai, Bassetto, Quarta & Mengali, 2019). The sensing, control, and learning algorithms that need to be embarked in such miniaturized devices can currently only be run on relatively heavy hardware, and need to be refined. However, nature has shown us that this is possible in the brains of insects with only a few hundred neurons. CA can also be decisive in other paradigms such as Nanotechnology, Ubiquitous Computing (López-de Ipiña, Chen, Mitton & Pan, 2017), and Quantum Computation (introduced by Feynman (Feynman, 1986)), particularly the Quantum CA (Lent, Tougaw, Porod & Bernstein, 1993; Watrous, 1995; Adamatzky, 2018). Due to the miniaturization hurdles of these devices, CA for stream learning may become of interest when there is no enough capacity to store huges volumes of data, and the computational capacity is very limited. After all, it is therefore not surprising that CA have received particular attention since the early days of CA investigation, and that CA for stream learning allows us to move in the correct direction in these scenarios, which are not far off the near future (Jafferis, Helbling, Karpelson & Wood, 2019).

2.2. Foundations of Cellular Automata

CA are usually described as discrete dynamical systems which present a universal computability capacity (Wolfram, 2018). Their beauty lies in its simple local interaction and computation of cells, which results in a huge complex behavior when these cells act together.

We can find four mutually interdependent parts in CA: i) the lattice, ii) its states,

iii) the neighborhood, and iv) the local rules. A *lattice* is created by a grid of elements (cells), which can be composed in one, two or higher dimensional space, but typically composed of uniform squared cells in two dimensions. CA contain a finite set of discrete states, whose number and range are dictated by the phenomenon under study. We find the simplest CA built using only one Boolean state in one dimension. The *neighborhood*, which is used to evaluate a *local rule*, is defined by a set of neighboring (adjacent) cells. A neighboring cell is any cell within a certain radius of the cell in question. Additionally, it is important to specify whether the radius R applies only along the axes of the cell space (von Neumann neighborhood), or if it can be applied diagonally (Moore neighborhood). In two dimensions, the R = 1 von Neumann neighborhood or the R = 1 Moore neighborhood are often selected (see Figure 1a). Finally, a *local rule* defines the evolution of each CA; it is usually realized by taking all states from all cells within the neighborhood, and by evaluating a set of logical or arithmetical operations written in the form of an algorithm (see Figure 1b). We can formally define a cellular automaton as follows, by adopting the notation used in (Kari, 2005): $A \doteq (d, S, f_{\boxplus}, f_{\bigcirc})$, where d represents the dimension, S a finite set of discrete states, $f_{\mathbb{H}}(\cdot)$ is a function that given a cell's coordinates at its input, returns the neighbors of the cell to be used in the update rule, and $f_{\odot}(\cdot)$ is a function that updates the state of the cell at hand as per the states of its neighboring cells. Therefore, in the case of a radius R = 1 von Neumann neighborhood defined over a d = 2-dimensional lattice, the set of neighboring cells and state of cell with coordinates c = [i, j] is given by:

$$f_{\mathbb{H}}([i,j]) = \{[i,j+1], [i-1,j], [i,j-1], [i+1,j]\},$$
(1)
$$S(\mathbf{c}) = S([i,j]) = f_{\mathcal{Q}}(S([i,j+1]), S([i-1,j]), S([i,j-1]), S([i+1,j])),$$
(2)

i.e., the states vector S([i, j]) of the [i, j] cell within the lattice is updated according to the local rule $f_Q(\cdot)$ applied over its neighbors given by $f_{\mathbb{H}}([i, j])$. In general, in a *d*-dimensional space, a cell's von Neumann neighborhood will contain 2d cells, and a Moore neighborhood will contain 3d - 1 cells. With this in mind, a cellular automaton should exhibit three properties to be treated as such: i) *parallelism* or *synchronicity* (all of the updates to the cells compounding the lattice are done at once); ii) *locality* (when a cell [i, j] is updated, its state S[i, j]) is based on the previous state of the cell and those of its nearest neighbors; and iii) *homogeneity* or *properties-uniformity* (the same update rule $f_Q(\cdot)$ is applied to each cell). The use of CA for pattern recognition is not straightforward. Some modifications and considerations should be performed before using CA for pattern recognition, since we need to map a dataset to a cell space:

• *Grid of cells* (lattice): despite being a natural fit to two-dimensional problems, CA must be extended to multiple dimensions to accommodate general pattern recognition tasks encompassing more dimensions. For *n* features, an approach adopted in the related literature is to assign one grid dimension to each feature of the dataset. Once dimensions of the grid have been set, we need to partition each grid dimension by the feature's values, obtaining an equal number of cells per dimension. To do that, evenly spaced values based on the maximum and minimum feature values (with an







Figure 1: Visual representation of CA for pattern recognition: (a) the von Neumann's (left) and Moore's (right) neighborhoods with radius R = 1; (b) the center cell examines its von Neumann's (left) and Moore's (right) neighborhoods and applies the local rule (majority vote) in a one-step update; (c) a two-dimensional dataset with instances $\mathbf{X}_t = (X_t^1, X_t^2)$ falling between [2,6] (min/max X_t^1) and [-2,-4] (min/max X_t^2), and a grid divided into G = 2 bins; (d) von Neumann's automata ($R = 1, d \times G$) ability to learn the data distribution from a few instances, and running up to 7 iterations of the initialization step (generations).

additional margin (*marg*) to ensure a minimal separation among feature's values) are used to create "bins" for each dimension of the data (see Figure 1c), which ultimately yield the cells of the lattice.

- *States*: a finite number of discrete states |S| are defined, corresponding to the number of classes considered in the task.
- Local rule: in pattern recognition tasks the update rule $f(\cdot)$ can be set to very assorted forms, being the most straightforward a majority voting among the states (labels) of its neighbors, i.e. for d = 2,

$$S([i,j]) = \underset{s \in \mathcal{S}}{\operatorname{arg\,max}} \sum_{[k,l] \in f_{\mathbb{H}}([i,j])} \mathbb{I}(S([k,l]) = s),$$
(3)

where $f_{\oplus}([i, j])$ returns the coordinates of neighboring cells of [i, j], and $\mathbb{I}(\cdot)$ is an auxiliary function taking value 1 if its argument is true (and 0 otherwise). Any other update rule can be defined to spread the state activation of each cell over its neighborhood (see Figure 1b).

- *Neighborhood*: it is necessary to specify a neighborhood and its radius. Although there are more types of local rules, "von Neumann" or "Moore" neighborhoods are often used (Figure 1a).
- *Initialization*: the grid is seeded depending on the feature values of the instances of the training dataset. Specifically, the state of each cell is assigned the label corresponding to the majority of training data instance with feature values falling within the range covered by the cell. As a result, cells will organize themselves into regions of similar labels (Figure 1d).
- *Generations*: after the initialization step, some cells can remain unassigned. Then, it is necessary to run the CA (generations) until no cells are left empty, and then continuing until either no changes are made or a fixed threshold is exceeded. Along this process, each cell computes its new state by applying the update rule over the cells in its immediate neighborhood. Each cell follows the same update rule, and all cells are updated simultaneously and synchronously. A critical characteristic of CA is that the update rule examines only its neighboring cells, so the processing is entirely local. No global or macro grid characteristics are computed whatsoever (Figure 1d).

2.3. Cellular Automata in Pattern Recognition

Despite the early findings presented in (Jen, 1986; Raghavan, 1993; Chaudhuri, Chowdhury, Nandi & Chattopadhyay, 1997), CA are not commonly used for pattern recognition. An exception is the work in (Fawcett, 2008), where CA were used as a form of instance-based classifiers for pattern recognition. These kinds of classifiers are well known by the pattern recognition community in the form of instance-based learning and nearest neighbors classifiers (Aha, Kibler & Albert, 1991; Duda, Hart & Stork,

2012). They represent regions of the instance space, so when a new instance needs to be classified, they select one or more close neighbors in these regions, and use their labels to assign a label to the new instance. Nevertheless, these are distinguished from CA in that they are not strictly local: there is no fixed neighborhood, hence an instance's nearest neighbor may change. In CA, by contrast, there is a fixed neighborhood, and the local interaction between cells influences the evolution and behavior of each cell. Fawcett, on the basis of Ultsch's work (Ultsch, 2002), put CA in value for the pattern recognition community by introducing them as a low-bias data mining method, simple but powerful for attaining massively fine-grained parallelism, non-parametric, with an effective and competitive classification performance in many cases (similar to that produced by other complex data mining models), and robust to noise. Besides, they were found to perform well with relatively scarce data. All this prior evidence makes CA suited for pattern recognition tasks.

Finally, it is worth highlighting the ability of CA to extract patterns from information and the possibility of generating tractable models, and being simple methods at the same time. This ability has encouraged the community to use them as a machine learning technique. Concretely, their traceability and reversibility (Kari, 2018) are renowned drivers for their adoption in contexts where model explainability is sought (Ribeiro, Singh & Guestrin, 2016; Gunning, 2017; Barredo Arrieta, Díaz-Rodrguez, Del Ser, Bennetot, Tabik, Barbado, García, Gil-López, Molina, Benjamins, Chatila & Herrera, 2019).

2.4. Cellular Automata for Stream Learning

In a real-time data mining process (stream learning), data streams are read and processed once per arriving sample (instance). Algorithms learning from such streams (stream learners) must operate under a set of constrained conditions (Domingos & Hulten, 2003):

- Each instance can be processed only once.
- The processing time of each instance must be low.
- Memory must be low as well, which implies that only a few instances of the stream should be explicitly stored.
- The algorithm must be prepared for providing an answer (i.e. a prediction) at any time of the process.
- Data streams evolve along time, which is an aspect that should be mandatorily considered.

In mathematical terms, a stream learning process evolving over time can be formally defined as follows: given a time period [0, t], we denote the historical set of instances as $\mathbf{D}_{0,t} = \mathbf{d}_0, \dots, \mathbf{d}_t$, where $\mathbf{d}_i = (\mathbf{X}_i, y_i)$ is a data instance, \mathbf{X}_i is the feature vector and y_i its label. We assume that $\mathbf{D}_{0,t}$ follows a certain joint probability distribution $P_t(\mathbf{X}, y)$. Such data streams are usually affected by non-stationary events (drifts) that eventually change their distribution (concept drift), making predictive models trained over these data obsolete. Bearing the previous notation in mind, concept drift at timestamp t + 1 occurs if $P_t(\mathbf{X}, y) \neq P_{t+1}(\mathbf{X}, y)$, i.e. as a change of the joint probability distribution of **X** and y at time t.

Since in stream learning we cannot explicitly store all past data to detect or quantify this change, concept drift detection and adaptation are acknowledged challenges for real-time processing algorithms (Lu, Liu, Dong, Gu, Gama & Zhang, 2018). Two strategies are usually followed to deal with concept drift:

- *Passive*, by which the model is continuously updated every time new data instances are received, ensuring a sufficient level of diversity in its captured knowledge to accommodate changes in their distribution; and
- Active, in which the model gets updated only when a drift is detected.

Both strategies can be successful in practice, however, the reason for choosing one strategy over the other is typically specific to the application. In general, a passive strategy has shown to be quite effective in prediction settings with gradual drifts and recurring concepts, while an active strategy works quite well in settings where the drift is abrupt. Besides, a passive strategy is generally better suited for batch learning, whereas an active strategy has been shown to work well in online settings (Gama, Medas, Castillo & Rodrigues, 2004; Bifet & Gavalda, 2007; Alippi, Boracchi & Roveri, 2013). In this work we have adopted an active strategy due to the fact that stream learning acts in an online manner.

Stream learning under non-stationary conditions is a field plenty of new challenges that require more attention due to its impact on the reliability of real-world applications. CA may provide a revolutionary view on RTA. Unfortunately, as shown later in Section 2.2, the original form of CA for pattern recognition does not allow for stream processing. A few timid attempts at incorporating CA concepts to RTA have been reported in the past. In (Hashemi, Yang, Pourkashani & Kangavari, 2007; Pourkashani & Kangavari, 2008) a cellular automaton-based approach was used as a real-time instance selector for stream learning. The classification task is carried out in batch mode by other non-CA-based learning algorithms. This is an essential difference with respect to the LUNAR approach proposed in this work, as the CA approach in (Hashemi, Yang, Pourkashani & Kangavari, 2007; Pourkashani & Kangavari, 2008) is not used for the learning task itself, but rather as a complement to the learning algorithm under use. Besides, they do not answer the relevant research questions posed in Section 1, which should be the basis for the use of CA in stream learning. The developed LUNAR algorithm is a streamified learning version of CA. It transforms CA into real incremental learners that incorporate an embedded mechanism for drift adaptation. In what follows we address the posed research questions by providing rationale on the design of LUNAR, and discussing on a set of experiments specifically devoted to inform our answers.

3. Proposed Approach: LUNAR

As it has been previously mentioned, LUNAR relies on the usage of CA for RTA. In this section we first introduce the modifications to *streamify* a CA pattern recognition approach (Subsection 3.1). Secondly, LUNAR is deeply detailed, grounded on the previously introduced material (Subsection 3.2).

3.1. Adapting Cellular Automata for Incremental Learning

The CA approach designed for pattern recognition can be adapted to cope with the constraints imposed by incremental learning. We present the details of this adaptation, which we coin as *streamified* Cellular Automaton (sCA), in Algorithm 1 (Streamified Cellular Automaton (sCA)):

- First, the sCA is created after setting the value of its parameters as per the dataset at hand (lines 1 to 5).
- Then, a set of *P* preparatory data are used to initialize the grid off-line by assigning the states of the corresponding cells according to the instances' values (lines 6 to 11).
- After this preliminary process, it is important to note that several preparatory data instances might collide into the same cell, thereby yielding several state occurrences for that specific cell. Since each cell must possess only one state for prediction purposes, lines 12 to 16 aim at removing this multiplicity of states by assigning each cell the state that occurs most among the preparatory data instances that fell within its boundaries.
- Similarly, we must ensure that all cells have a state assigned, i.e. no empty grid cell status must be guaranteed. To this end, lines 17 to 23 apply the local rule $f_Q(\cdot)$ over the neighbors of every cell, repeating the process (generations) until every cell has been assigned a state.
- This procedure can again render several different states for a given cell, so the previously explained majority state assignment procedure is again enforced over all cells of the grid (lines 24 to 28).

Once this preparatory process is finished, the sCA is ready for the *test-then-train* (Gama, Žliobaitė, Bifet, Pechenizkiy & Bouchachia, 2014) process with the rest of the instances (lines 29 to 34). In this process, the sCA first predicts the label of the arriving instance (testing phase), and updates the limits to reconfigure the bins (training phase). By this way, the sCA always represents the currently prevailing distribution of the streaming data at its input. After that, the sCA updates the current state of the cell enclosing the instance with the true label of the incoming instance. As a result, the cells of sCA always have an updated state according to the streaming data distribution.

The details of the proposed LUNAR algorithm underline the main differences between the CA version for pattern recognition and sCA. We highlight here the most relevant ones:

Algorithm 1: Streamified Cellular Automaton (sCA)

```
Input : Preparatory data instances [(\mathbf{X}_t, y_t)]_{t=0}^{t=P-1}; training/testing data for the
                 rest of the stream [(\mathbf{X}_t, y_t)]_{t=P}^{\infty}; the grid size G (bins per dimension); a
                 local update rule f_{\bigcirc}(\cdot); a neighborhood function f_{\boxplus}(\mathbf{c}) for cell with
                 coordinates \mathbf{c} \in \mathcal{G} = \{1, \dots, G\}^d as its argument; a radius R for the
                 neighborhood operator
    Output: Trained streamified CA (sCA), producing predictions \hat{y}_t \ \forall t \in [P, \infty)
 1 Let the number of dimensions d of the grid be the number of features in X_t
 2 Let the number of cell states |\mathcal{S}| be the number of classes (alphabet of y_t)
 3 Set an empty vector of state hits per every cell: \mathbf{h}_{\mathbf{c}} = \begin{bmatrix} 1 & \forall \mathbf{c} \in \mathcal{G} \end{bmatrix}
 4 Initialize the limits of the grid: [(lim_n^{low}, lim_n^{high})]_{n=1}^d
 4 Initialize the limits of the grid: [(lim_n^{low}, lim_n^{ness})]_{n=1}^{a}

5 Create the sCA grid as per G, n and [(lim_n^{low}, lim_n^{high})]_{n=1}^{d}
 6 for t = 0 to P - 1 do // Preparatory process
         Update limits as per \mathbf{X}_t, e.g., lim_n^{low} = \min\{lim_n^{low}, x_t^n\}
 7
         Reconfigure grid bins as per G and the updated [(lim_n^{low}, lim_n^{high})]_{n=1}^d
 8
         Select the cell c in the grid that encloses X_t
 9
         Append y_t to the vector of state hits in the cell, e.g. \mathbf{h}_{\mathbf{c}'} = [\mathbf{h}_{\mathbf{c}'}, y_t]
10
11 end
12 for \mathbf{c} \in \mathcal{G} do // Guaranteeing one state per cell
         if |\mathbf{h_c}| > 1 then // More than 1 state hit in cell
13
          \int S(\mathbf{c}) = \arg \max_{s \in \mathcal{S}} \sum_{i=1}^{|\mathbf{h}_{\mathbf{c}}|} \mathbb{I}(h_{\mathbf{c}}^{i} = s)
14
         end
15
16 end
    while \exists \mathbf{c} \in \mathcal{G} : h_{\mathbf{c}} = [] \mathbf{do} / / Ensuring a hit in all cells
17
         for c \in \mathcal{G} do
18
              Compute the neighboring cells of c as per f_{\oplus}(c) and R
19
              Compute state s' by applying rule f_{\bigcirc}(\cdot) over the neighboring cells
20
              Append s' to h_{\mathbf{c}}, i.e., \mathbf{h}_{\mathbf{c}} = [\mathbf{h}_{\mathbf{c}}, s']
21
         end
22
23 end
24 for \mathbf{c} \in \mathcal{G} do // Guaranteeing one state per cell
         if |\mathbf{h_c}| > 1 \text{ then } // More than 1 state hit in cell
25
          S(\mathbf{c}) = \arg \max_{s \in \mathcal{S}} \sum_{i=1}^{|\mathbf{h}_{\mathbf{c}}|} \mathbb{I}(h_{\mathbf{c}}^{i} = s)
26
         end
27
28 end
29 for t = P to \infty do // Stream learning
         Predict \hat{y}_t as S(\mathbf{c}), with \mathbf{c} denoting the coordinates of the cell enclosing \mathbf{X}_t
30
         Update limits as per \mathbf{X}_t, e.g., lim_n^{low} = \min\{lim_n^{low}, x_t^n\}
31
         Reconfigure grid bins as per G and the updated [(lim_n^{low}, lim_n^{high})]_{n=1}^d
32
         Update cell state to the verified class of the test instance: S(\mathbf{c}) = y_t
33
34 end
```

- *Preparatory instances*: a small portion of the data stream (preparatory instances $[(\mathbf{X}_t, y_t)]_{t=0}^{P-1}$) is used to perform the sCA initialization. Then, the grid is seeded with these data values, and sCA progresses through several generations (and applying the local rule) until all cells are assigned a state. With the traditional CA for pattern recognition, the whole dataset is available from the beginning. Therefore, the CA initialization is not required.
- An updated representation of the instance space: since historical data is not available and data grow continuously, the sCA updates the cell bins of the grid every time a new data instance arrives. By doing this, we ensure that the range of values of the instance space is updated during the streaming process. With the traditional CA for pattern recognition, as the whole training dataset is static, the grid boundaries and cell bins are calculated at the beginning of the data mining process, and remain unaltered during the test phase.
- *CA's cells continuously updated*: sCA's cells are also updated when new data arrives by assigning the incoming label (state) to the corresponding cell. Instead of examining the cells in the immediate neighborhood of this corresponding cell during several generations (which would take time and computational resources), sCA just checks the previous state of the cell and the current state provided by the recent data instance. When both coincide, no change is required, but when they differ, the cell adopts the state of the incoming data instance. In this sense, sCA always represents the current data distribution. This is the way sCA works as an incremental learner. However, with the traditional CA for pattern recognition, once initialized, a local rule is applied over successive generations to set the state of a given cell depending on the states of its neighbors.

3.2. LUNAR: a sCA with Drift Detection and Adaptation Abilities

LUNAR blends together sCA and paired learning in order to cope with concept drift. In essence, a stable sCA is paired with a reactive one. The scheme inspires from the findings reported in (Bach & Maloof, 2008): a stable learner is incrementally trained and predicts considering all of its experience during the streaming process, whereas a reactive learner is trained and predicts based on its experience over a time window of length W. Any online learning algorithm can be used as a base learner for this strategy. As shown in Algorithm 2 (Paired learning for OLMs), the stable learner is used for prediction, whereas the reactive learner is used to detect possible concept drifts:

- While the concept remains unchanged, the stable learner provides a better performance than the reactive learner (lines 6 and 7).
- A counter is arranged to count the number of times within the window that the reactive learner predicts correctly the stream data, and the stable learner produces a bad prediction (lines 8 to 10).

- If the proportion of these occurrences surpasses a threshold θ (line 11), a concept change is assumed to have occurred, the stable learner is replaced with the reactive one, and the stable learner is reinitialized.
- Finally, both stable and reactive learners are trained incrementally (lines 15 and 16).

Algorithm 2: Paired learning for OLMs **Input** : Preparatory data instances $\{(\mathbf{X}_t, y_t)\}_{t=0}^{t=P-1}$; training/testing data for the rest of the stream $\{(\mathbf{X}_t, y_t)\}_{t=P}^{\infty}$; window size W for the reactive learner; threshold θ for substituting the stable learner with the reactive one; stream learning model $f(\mathbf{X})$ **Output:** Updated stable and reactive learners $f_s(\cdot)$ and $f_r(\cdot)$ 1 Let $\mathbf{c} = [c^w]_{w=1}^W$ be a circular list of W bits, each initially set to $c^w = 0 \ \forall w$ **2** for t = 0 to P - 1 do // Preparatory process Train both $f_s(\cdot)$ and $f_r(\cdot)$ with (\mathbf{X}_t, y_t) 3 4 end **5** for t = P to ∞ do // Stream learning Let $\widehat{y}_s = f_s(\mathbf{X}_t)$ (output of the model) and $\widehat{y}_r = f_r(\mathbf{X}_t)$ 6 Set $c^1 = 0$ and $c^w = c^{w-1}$ for w = 2, ..., W7 **if** $\widehat{y}_s \neq y_t$ and $\widehat{y}_r = y_t$ then 8 Set $c^1 = 1$ 9 10 end if $W^{-1}\sum_{w=1}^{W} c^w > \theta$ then // Drift detection 11 Replace $f_s(\cdot)$ with $f_r(\cdot)$ (including its captured knowledge) 12 Set $c^w = 0 \ \forall w = 1, \dots, W$ 13 14 end Train $f_s()$ incrementally with (\mathbf{X}_t, y_t) 15 Train $f_r(\cdot)$ from scratch over $\{(\mathbf{X}_{t'}, y_{t'})\}_{t'=t-W+1}^t$ 16 17 end

Algorithm 3 (LUNAR) describes in detail the proposed algorithm. In essence the overall scheme relies on a couple of paired sCA learners: one in charge of making predictions on instances arriving in the data stream (*stable* sCA learner), whereas the other (*reactive* sCA learner) captures the most recent past knowledge in the stream (as reflected by a time window W). As previously explained for generic paired learning, the comparison between the accuracies scored by both learners over the time window permits to declare the presence of a concept drift when the reactive learner starts predicting more accurately than its stable counterpart. When this is the case, the grid state values of the reactive sCA is transferred to the stable automaton, and proceeds forward by predicting subsequent stream instances with an updated grid that potentially best represents the prevailing distribution of the stream.

We now summarize the most interesting capabilities of the LUNAR algorithm:

• *Incremental learning*: as it relies on sCA learners, LUNAR shows a natural ability to learn incrementally, which is a crucial requirement under the constrained conditions

Algorithm 3: LUNAR

```
Input : Preparatory data instances \{(\mathbf{X}_t, y_t)\}_{t=0}^{P-1}; training/testing data for the
                rest of the stream (\mathbf{X}_t, y_t)_{t=P}^{\infty}; window size W for the reactive sCA
                learner; threshold \theta for substituting the stable sCA learner with the
                reactive one; sCA parameters G, f_{\bigcirc}(\cdot), f_{\boxplus}(\mathbf{c}) and R as per Algorithm
   Output: Updated stable and reactive learners sCA_s and sCA_r
 1 Let \mathbf{\hat{c}} = [c^w]_{w=1}^W be a circular list of W bits, each initially set to c^w = 0 \ \forall w
 2 Perform lines 1 – 28 of Algorithm 1 for sCA<sub>s</sub> and sCA<sub>r</sub> over \{(\mathbf{X}_t, y_t)\}_{t=0}^{P-1}
 3 for t = P to \infty do // Stream learning
        Let \widehat{y}_s = \text{sCA}_s(\mathbf{X}_t) and \widehat{y}_r = \text{sCA}_r(\mathbf{X}_t) as per line 30 of Algorithm 1
 4
        Set c^1 = 0 and c^w = c^{w-1} for w = 2, ..., W
 5
        if \widehat{y}_s \neq y_t and \widehat{y}_r = y_t then
 6
           Set c^1 = 1
 7
        end
 8
        if W^{-1}\sum_{w=1}^{W} c^w > \theta then // Drift detection
 9
             Copy grid state values of sCA_r to sCA_s (knowledge transfer)
10
             Clear sCA<sub>r</sub> and seed it with \{(\mathbf{X}_{t'}, y_{t'})\}_{t'=t-W+1}^t
11
             Perform lines 1 - 28 of Algorithm 1 for sCA_r
12
             Set c^w = 0 \ \forall w = 1, \dots, W
13
        end
14
        Train sCA<sub>s</sub> incrementally with (\mathbf{X}_t, y_t) // Lines 31-33, Alg.1
15
        Train sCA<sub>r</sub> from scratch over \{(\mathbf{X}_{t'}, y_{t'})\}_{t'=t-W+1}^{t}
16
17 end
```

present in RTA.

- *Few parameters*: LUNAR requires only a few parameters, a warmly welcome characteristic that simplifies the parameter tuning process in drifting environments.
- Little data to be trained: LUNAR can learn a data distribution with relatively few instances which, in the context of stream learning, involves a quicker preparation (warm-up) phase. However, depending on the grid size G, several generations may be required to make all cells be assigned a state. This is exemplified in Figure 1d, which considers a dataset with 2 continuous features $\mathbf{X}_t = (X_t^1, X_t^2) \in \mathbb{R}[0, 1]$, and a binary target class $y_t \in \{0, 1\}$. Here, a $d \times G = 2 \times 5$ cellular automaton only needs 2 iterations of the loops in lines 12 to 28 (hereafter referred to as generation) until all cells are assigned one and only one state (see Algorithm 1). However, the grid is coarse, and its distribution of states does not represent the distribution of the data stream. With a $d \times G = 2 \times 10$ cellular automaton, 4 generations are needed until no cells are left empty, and the representation of the data distribution is more finely grained. Finally, the $d \times G = 2 \times 20$ cellular automaton needs 7 generations until

no cells are left empty, yet the representation of the data distribution is well defined. A good balance between representativeness and complexity must be met in practice when deploying LUNAR in real setups.

- *Constrained scenarios*: in setups where computational costs must be reduced even more, a subsampling strategy is often recommended, where only a fraction of instances of the data stream is considered for model updating. In these scenarios, the capability of CA to represent the data distribution with a few instances fits perfectly. So does LUNAR by embracing sCA at its core.
- Stream learning under verification latency: learning in an environment where labels do not become immediately available for model updating is known as verification latency. This requires mechanisms and techniques to propagate class information forward through several time steps of unlabeled data. This remains as an open challenge in the stream learning community (Krempl, Žliobaite, Brzeziński, Hüllermeier, Last, Lemaire, Noack, Shaker, Sievi, Spiliopoulou & Stefanowski, 2014). Here, LUNAR finds its gap as well, due to its capability of evolving through several generations and represents a data distribution from a few annotated data instances.
- *Evolving scenarios*: by harnessing the concept of paired learners, LUNAR adapts to evolving conditions where changes (drifts) provoke that learning algorithms have to forget and learn the old and the new concept, respectively.
- *Tractable and interpretable*: both characteristics lately targeted with particularly interest under the eXplainable Artificial Intelligence (XAI) paradigm (Barredo Arrieta, Díaz-Rodrguez, Del Ser, Bennetot, Tabik, Barbado, García, Gil-López, Molina, Benjamins, Chatila & Herrera, 2019).

Finally, LUNAR enters the stream learning scene as a new base learner for the stateof-the-art.

4. Experimental Setup

We have designed two experiments in order to answer the three research questions posed in the Section 1. The first experiment addresses the research questions RQ1 and RQ2 on the capability of sCA to learn incrementally and adapt to changes conditions, while the second experiment addresses RQ3 by discussing on the competitiveness of our LUNAR algorithm with respect to other streaming learners.

In all the experiments we have used the von Neumann neighborhood because it is linear in the number of dimensions of the instance space, so it scales well when dealing with problems of high dimensionality. In addition, this neighborhood is composed of less neighbors than in Moore's case, and the local rule is applied over less cells, which makes the process lighter and better for streaming scenarios in terms of computational and processing time costs. Regarding the performance measurement, we have adopted the so-called *prequential accuracy* (Dawid, Vovk et al., 1999) for all experiments, as it is a suitable metric to evaluate the learner performance in the presence of concept drift (Dawid, Vovk et al., 1999; Gama, Sebastião & Rodrigues, 2013; Gama, Žliobaitė, Bifet, Pechenizkiy & Bouchachia, 2014). This metric quantifies the average accuracy obtained by the prediction of each test instance before its learning in an online *test-then-train* fashion, and is defined as:

$$preACC(t) = \begin{cases} preACC_{ex}(t), & \text{if } t = t_{ref}, \\ preACC_{ex}(t-1) + \frac{preACC_{ex}(t) - preACC_{ex}(t-1)}{t - t_{ref} + 1}, & \text{otherwise,} \end{cases}$$
(4)

where $preACC_{ex}(t) = 0$ if the prediction of the test instance at time t before its learning is wrong and 1 if it is correct; and t_{ref} is a reference time that fixes the first time step used in the calculation. This reference time allows isolating the computation of the prequential accuracy before and after a drift has started.

As for the initialization of sCA, following Algorithm 1 we have used a reduced group of P preparatory instances. In the second experiment we have also used these instances to carry out the hyper-parameter tuning of the OLMs under analysis. The selection of the size of this preparatory data usually depends on the available memory or the processing time we can take to collect or process these data. Finally, following the recommendations of (Fawcett, 2008), we have assigned one grid dimension to each attribute.

4.1. First Experiment: Addressing RQ1 and RQ2 with sCA

In order to address RQ1 and RQ2, we have used several two-dimensional synthetic datasets for the first experiment, where results can be easily visualised and interpreted. Synthetic data are advisable because in real datasets is not possible to know exactly when a drift appears, which type of drift emerges, or even if there is any confirmed drift. Thus, it is not possible to analyze the behavior of sCA in the presence of concept drift by only relying on real-world datasets.

This being said, for this experiment we have used the sCA detailed in Section 3.1 (Algorithm 1). We have compared its naive version with the one using an active adaptation mechanism: assuming a perfect drift detection (the drift point is known beforehand), once a drift occurs the grid of the sCA model is seeded with a *W*-sized window of past instances, which allows representing the current concept (data distribution). Then, sCA progresses again through several generations (and applying the local rule) until all cells are assigned a state. As highlighted before, this ability of CA to represent a data distribution from a few instances is very valuable here, where only a limited sliding window is used to initialize the sCA grid. This experiment is designed to illustrate and visualize the operation of sCA in two dimensions (two features).

We use the renowned set of four balanced synthetic datasets (CIRCLE, LINE, SINEH, and SINEV) described in (Minku, White & Yao, 2009). Very briefly, these datasets contain one simulated drift characterized by low and high speed, resulting 2 different types of drift for each dataset. Speed is the inverse of the time taken for a

new concept to completely replace the previous one. Each dataset has 2,000 instances $(t \in \{1, ..., 2, 000\})$, 2 normalized ([0,1]) continuous features $\mathbf{X}_t = (X_t^1, X_t^2)$, and a binary target class $y_t \in \{0, 1\}$. Drift occurs at t = 1,000 (where it goes from the old concept to the new one), and the drifting period is 1 for abrupt drifts (high speed datasets) and 500 for gradual ones (low speed datasets). For abrupt drifts we have opted for a small window size of W = 25, whereas for gradual drifts we need a bigger window of W = 100 (Gama, Sebastião & Rodrigues, 2013). The size of this window will depend on the type of drift: a small window can assure fast adaptability in abrupt changes, while a large window produces lower variance estimators in stable phases, but cannot react quickly to gradual changes. Thus, sliding windows is a very common form of the so-called *forgetting mechanism* (Gama, 2010), where outdated data is discarded and adapt to the most recent state of the nature, then emphasis is placed on error calculation from the most recent data instances.

4.2. Second Experiment: Addressing RQ3 with LUNAR

For the second experiment we have resorted to real-world datasets to confirm the competitive performance of LUNAR in a comparison with recognized OLMs in the literature, answering RQ3. In this case, since we deal with problems having n features, an extensive use is to assign one grid dimension of the CA to each feature of the dataset. In this experiment, LUNAR is compared with paired learning models (see Algorithm 2) using at their core different online learning algorithms from the literature. These comparison counterparts have been selected over others due to the fact that they are well-established methods in the stream learning community. Also because their implementations are reliable and easily accessible in well-known Python frameworks such as scikit-multiflow (Montiel, Read, Bifet & Abdessalem, 2018) and scikit-learn (Pedregosa, Varoquaux, Gramfort, Michel, Thirion, Grisel, Blondel, Prettenhofer, Weiss, Dubourg et al., 2011). As this is the first experiment carried out to know about their performance as incremental learners with drift adaptation, we have selected these reputed methods:

- *Stochastic Gradient Descent Classifier* (SGDC) (Bottou, 2010) implements an stochastic gradient descent learning algorithm which supports different loss functions and penalties for classification.
- *Hoeffding Tree* (HTC) (Domingos & Hulten, 2000), also known as Very Fast Decision Tree (VFDT), is an incremental anytime decision tree induction algorithm capable of learning from massive data streams, assuming that the distribution of data instances does not change over time, and exploiting the fact that a small instance can often be enough to choose an optimal splitting attribute.
- Passive Agressive Classifier (PAC) (Crammer, Dekel, Keshet, Shalev-Shwartz & Singer, 2006) focuses on the target variable of linear regression functions, $\hat{y}_t = \boldsymbol{\omega}_t^T \cdot \mathbf{X}_t$, where $\boldsymbol{\omega}_t$ is the incrementally learned weight vector. After a prediction is made, the algorithm receives the true target value y_t and an instantaneous ε -insensitive hinge

loss function is computed to update the weight vector. This loss function was specifically designed to work with stream data, and is analogous to a standard hinge loss. The role of ε is to allow for a lower tolerance of prediction errors. Then, when a round finalizes, the algorithm uses ω_t and the instance (\mathbf{X}_t, y_t) to produce a new weight vector ω_{t+1} , which is then used to predict the next incoming instance.

• *k-Nearest Neighbours* (KNN) (Read, Bifet, Pfahringer & Holmes, 2012), a wellknown lazy learner where the output is given by the labels of the K training instances closest (under a certain distance measure) to the query instance X_t . In its *streamified* version, it works by keeping track of a sliding window of past training instances. Whenever a query request is executed, the algorithm will search over its stored instances and find the $K \leq W$ closest neighbors in the window, again as per the selected distance metric.

Three real-world datasets have been utilized in this second experiment:

- The Australian New South Wales Electricity Market dataset (ELEC2) (Gama, Medas, Castillo & Rodrigues, 2004) is a widely adopted real-world dataset in studies related to stream processing/mining, particularly in those focused on non-stationary settings. In essence the dataset represents electricity prices over time, which are not fixed and become affected by the demand and supply dynamics. It contains 45,312 instances dated from 7 May 1996 to 5 December 1998. Each instance X_t of the dataset refers to a period of 30 minutes, and has d = 5 features (day of week, time stamp, NSW electricity demand, Vic electricity demand, and scheduled electricity transfer between states). The target variable y_t to be predicted is binary, and identifies the change of the price related to a moving average of the last 24 hours. The class level only reflect deviations of the price on a one day average, and removes the impact of longer term price trends.
- The Give Me Some Credit dataset (GMSC¹) correspond to a credit scoring task in which the goal is to decide whether a loan should be granted or not (binary target y_t). This is a core decision for banks due to the risk of unexpected expenses and future lawsuits. The dataset comprises supervised historical data of 150,000 borrowers described by d = 10 features.
- The POKER-HAND dataset² originally consists of 1,000,000 stream instances. Each record is an example of a hand consisting of five playing cards drawn from a standard deck of 52 cards. Each card is described by 2 attributes (suit and rank), giving rise to a total of d = 10 predictive attributes for every stream instance X_t . The goal is

¹Available at: https://www.kaggle.com/c/GiveMeSomeCredit. Last accessed on December 5th, 2019.

²Available at: https://archive.ics.uci.edu/ml/datasets/Poker+Hand. Last accessed on December 5th, 2019.

to predict the poker hand upon such features, out of a set of |S| = 10 possible poker hands (i.e., multiclass classification task).

In order to alleviate the computational effort needed to run the entire benchmark, we have selected 20,000 instances of each dataset. We have considered the first 50% of the datasets (10,000 instances) to tune the parameters of the OLMs and to seed LUNAR, whereas the rest 50% has been used for prediction and performance assessment. Each experiment has been carried out 25 times in the case of SGDC, PAC and KNNC to account for the stochasticity of the hyperparameter search algorithm and their learning procedure. Indeed, the considered online models are sensible to parameter values. Thus we have performed a randomized hyper-parameter tuning over preparatory instances, obtaining different hyper parameter settings for each run. Table 1 summarizes the parameters that have participated in each randomized search process, following the nomenclature of the scikit-learn³ and scikit-multiflow⁴ frameworks. In the case of HTC, we found experimentally that its default values (see documentation of scikit-multiflow for more details) have worked very well in all datasets, so it was not necessary to carry out the randomized search on hyper parameters, which helped speed up the experimentation. Finally, the same window size W than LUNAR has been assigned to KNNC for the sake of a fair comparison between them.

Model	Parameters	Values used	Tuning
SGDC	alpha loss learning_rate eta0 penalty max_iter	$\begin{array}{c} 10^{-x}, x \in \{-1,-2,\ldots,-6\}\\ \text{`perceptron', `hinge', `log', `modified_huber', `squared_hinge'}\\ \text{`constant', `optimal', `invscaling', `adaptive'}\\ 0.1, 0.5, 1.0\\ \text{None, `l2', `l1', `elasticnet'}\\ 1, 100, 200, 500\\ \end{array}$	yes
PAC	C max_iter	$\begin{array}{c} 0.001, 0.005, 0.01, 0.05, 0.1, 0.5, 1.0 \\ 1, 100, 200, 500 \end{array}$	yes
KNNC	n_neighbors leaf_size algorithm weights	5, 10, 15, 25, 50 5, 10, 20, 30 'auto' 'uniform', 'distance'	yes
HTC	grace_period split_criterion leaf_prediction nb_threshold	200 ʻinfo_gain' ʻnba' 0	no

Table 1: Parameters that have participated in each randomized searching process of SGDC, PAC and KNNC. In the case of HTC its default values show a good enough performance and has not participated in the randomized hyperparameter searching process. This has alleviated the computing cost of the experiments.

Finally, Table 2 summarizes the parameter settings of LUNAR and the OLMs used for the real-world experiments.

³https://scikit-learn.org/. Last access in December 5th, 2019.

⁴https://scikit-multiflow.github.io/. Last access in December 5th, 2019.

Model	Parameter		Real data	Synthetic data					
		ELEC2	GMSC	POKER-HAND	~ 5				
All models	Р	50% of t	he considered st	5% of the considered stream length					
	W	50 instances	250 instances	250 instances	25, 50, 100 instances				
	$f_{\boxplus}(\cdot)$	von Neumann							
Ц	$f_{Q}(\cdot)$								
NA	R	1 (�)							
ΓΩ	S	$\{0, 1\}$	$\{0, 1\}$	$\{0,1,\ldots,9\}$	$\{0,1\}$				
	$d \times G$	5×5	10×3	10×3	$2 \times 5, 2 \times 10, 2 \times 20$				
	θ	0.05	0.01	0.001	Not applicable				
SGDC		0.8	0.001	0.1					
HTC		0.1	0.0001	0.01					
PAC	θ	0.8	0.001	0.1	Not used				
KNNC		0.1	0.001	0.001					

Table 2: Parameters configuration for LUNARand the considered OLMs in synthetic (RQ1 and RQ2) and real-world experiments (RQ3).

5. Results and Analysis

Next, we present the results for the two experiments, organizing the foregoing analysis and discussion in terms of the research questions posed in Section 1:

5.1. RQ1: Does sCA act as a real incremental learner?

Definitely yes. If we combining the definition of incremental learning suggested by (Giraud-Carrier, 2000; Lange & Zilles, 2003) with Kuncheva's and Bifet's desiderata for non-stationary learning (Kuncheva, 2004; Bifet, Gavaldà, Holmes & Pfahringer, 2018), incremental learning can be understood as the capacity to process and learn from data in an incremental fashion, and to deal with data changes (drifts) that may occur in the environment. Therefore, the properties that guide the creation of incremental learning algorithms (Ditzler & Polikar, 2012) are:

- Learning new knowledge: this feature is essential for online learners that have to be deployed over non-stationary settings. To provide evidence that SCA incorporates this characteristic, Figures 2 and 4 summarize the performance results obtained over two synthetic datasets (CIRCLE and SINEH) with an abrupt drift occurring at t = 1000, and assuming to have been detected at t = 1025. The adaptation mechanism resorts to a window of W = 25 past data instances; upon the detection of the drift, the entire knowledge contained in the cellular automata (i.e. the distribution of cell states) is erased, and the automata is seeded with the instances falling in the last W-sized window. Although the abrupt drift obliges to learn quickly the new concept and forgetting the old one, we observe in these plots that sCA excels at this task, yielding a prequential accuracy over time that does not get apparently affected by the change of concept.
- *Preserving previous knowledge*: related to the previous property, gradual drifts require maintaining part of the previous concept over time, so that the evolved model

leverages the retained knowledge during its transition to the new concept. This is clearly shown in Figures 3 and 5, which both focus on CIRCLE and SINEH respectively, showing a gradual drift at t = 1000, and assuming a drift detection at t = 1600. The adaptation mechanism resorts to a window of W = 100 past data instances. In particular, the prequential accuracy results and subplots therein included show that sCA can preserve the old concept while learning the new one through the implementation of a forgetting strategy.

- One-pass (incremental) learning: through preceding sections we have elaborated on the incremental learning strategy of sCA (lines 30 to 33 in Algorithm 1), by which it is able to learn one instance at a time without requiring any access to historical data.
- *Dealing with concept drift*: Finally, we have seen in the previous experiments that sCA is capable of dealing with evolving conditions by using a drift detection and by implementing an adaptation scheme. We will later revolve around these results.



Figure 2: Learning and adaptation of von Neumann's sCA of different grid sizes for the CIRCLE dataset, which exhibits an abrupt drift at t = 1000. The drift is detected at t = 1025 and then the adaptation mechanism, which uses a window of instances W = 25, is triggered. (a) Performance comparison at points **b** (initial), **c** (just before the drift occurs), **d** (W = 25 instances after drift detection and the adaptation mechanism were triggered), and **e** (final). The dashdotted line points out the drift detection and the dotted line the point in which the performance is measured. The shaded area corresponds to the preparatory instances. (b) The cells of sCAs are seeded with the preparatory instances (**b**). (c) The learning process of sCA before drift occurs (**c**). (d) The learning process of sCA after the drift occurs, and how those with the adaptation mechanism is initialized and seeded with a set of W = 25 past instances (**d**). (e) The learning process of sCA until the end of the stream (**e**).



Figure 3: Learning and adaptation of von Neumann's sCA of different grid sizes for the CIRCLE dataset, which exhibits a gradual drift at t = 1000. The drift is detected at t = 1600, and then the adaptation mechanism, which uses a window of instances W = 100, is triggered. The interpretation of the remaining plots is the same than those in Figure 2.



Figure 4: Learning and adaptation of von Neumann's sCA of different grid sizes for the SINEH dataset, which exhibits an abrupt drift at t = 1000. The drift is detected at t = 1025 and then the adaptation mechanism, which uses a window of instances W = 25, is triggered. The interpretation of the remaining plots is the same than those in Figure 2.



Figure 5: Learning and adaptation of von Neumann's sCA of different grid sizes for the SINEH dataset, which exhibits a gradual drift at t = 1000. The drift is detected at t = 1600 and then the adaptation mechanism, which uses a window of instances W = 100, is triggered. The interpretation of the remaining plots is the same than those in Figure 2.

5.2. RQ2: Can sCA successfully adapt to evolving conditions?

In a nutshell, the answer is yes. However, the success rate depends on the type of drift and the size G of the CA grid.

This is empirically shown in Figures 2 and 4 for abrupt drifts, where we have applied an adaptation mechanism to a sCA of different grid sizes over the CIRCLE and SINEH datasets, respectively. Once the drift appears at t = 1000 (solid vertical line) and is detected (dashdotted vertical line), we observe that sCA with and without adaptation mechanism perform equal at points **b** and **c**. But when the adaptation mechanism is triggered we see at point d (25 instances after drift detection and adaptation) and e (the end of the data stream) that sCA using grids of $d \times G = 2 \times 10$ and $d \times G = 2 \times 20$ obtain better prequential accuracies when incorporating an adaptation mechanism. In the case of CIRCLE dataset, the adaptive versions of sCA obtain an average prequential accuracy of $0.873 (2 \times 10)$ and $0.900 (2 \times 20)$ versus their non-adaptive counterparts, which respectively score accuracies of 0.866 and 0.880. The same situation holds with the SINEH dataset, where again adaptive sCAs obtain average prequential accuracies of $0.773 (2 \times 10)$ and $0.833 (2 \times 20)$ against their non-adaptive versions (0.765 and 0.808, respectively). The adaptive versions are able to forget quickly the old concept because they are reinitialized with a few instances (25) of the new concept, as opposed to the non-adaptive versions which keep the old concept along time (see Figures 2d and 4d). At point e (Figures 2e and 4e), we can observe some differences in the learning of both versions. In the case of the small grid size 2×5 , neither adaptive nor non-adaptive versions are able to learn the concepts properly and establishing the boundaries between classes, yielding a poor classification performance. Therefore, when the change from the old to the new concept is abrupt, the fast forgetting of the old concept becomes primordial (Gama, Sebastião & Rodrigues, 2013). For a complete comparison with more datasets we refer to Table 3.

Now, we focus on gradual drifts of CIRCLE and SINEH dataset (Figures 3 and 5 respectively), where a gradual drift also occurs at (t = 1,000). Figures 3a and 5a, and Table 3, show how both sCA with and without adaptation mechanism obtain almost the same mean prequential accuracy. As the old concept disappears slowly while the new one also does it slowly, the adaptive version with a window of instances (w = 100) and the non-adaptive version do preserve the old concept while learn the new one. Even so, we can observe slight improvements of adaptive versions over the non-adaptive ones at points **d** and **e**. These differences are bigger in the case of sCA with a 2×20 grid: 0.894 for the adaptive version against 0.888 for the non-adaptive version (CIRCLE at point d), 0.897 for the adaptive version against 0.881 for the non-adaptive version (CIRCLE at point e), 0.812 for the adaptive version against 0.811 for the non-adaptive version (SINEH at point d), and 0.820 for the adaptive version against 0.811 for the non-adaptive version (SINEH at point e). Again, in the case of a 2×5 grid, due to the small grid size neither adaptive nor non-adaptive versions are able to learn the concepts properly and establishing the boundaries between classes, achieving a poor classification performance. For the case of $sCA2 \times 10$, because the drift is gradual, the grid size does not allow for capturing well enough the differences of both

concepts. Finally, when the change from the old concept to the new one is gradual, an slow forgetting of the old concept becomes primordial (Gama, Sebastião & Rodrigues, 2013).

Synthetic	sCA	$\mathbf{d}\times \mathbf{G}$	$\mathbf{preACC}(\mathbf{t})$			Synthetic	sC2	d×G	$\mathbf{preACC}(\mathbf{t})$				
datasets			С	d	e	Average	datasets	asets	und	С	d	e	Average
CIRCLE (abrupt)	Adaptive	2×5	0.827	0.822	0.827	0.794		Adaptive	2×5	0.833	0.822	0.811	0.777
		2×10	0.928	0.918	0.905	0.873			2×10	0.933	0.920	0.921	0.877
		2×20	0.958	0.943	0.931	0.900	SINEV		2×20	0.960	0.947	0.948	0.907
	Non-adaptive	2×5	0.827	0.822	0.828	0.795	(abrupt)	(abrupt) Non-adaptive	2×5	0.833	0.823	0.811	0.777
		2×10	0.928	0.915	0.892	0.866			2×10	0.933	0.912	0.905	0.867
		2×20	0.958	0.939	0.881	0.880			2×20	0.960	0.939	0.876	0.879
		2×5	0.827	0.816	0.820	0.789		Adaptive	2×5	0.833	0.753	0.756	0.757
	Adaptive	2×10	0.928	0.885	0.890	0.867			2×10	0.933	0.881	0.891	0.866
CIRCLE (gradual)		2×20	0.958	0.894	0.897	0.890	SINEV		2×20	0.960	0.893	0.906	0.893
	Non-adaptive	2×5	0.827	0.817	0.820	0.789	(gradual)		2×5	0.833	0.754	0.757	0.757
		2×10	0.928	0.886	0.891	0.868		Non-adaptive	2×10	0.933	0.882	0.892	0.867
		2×20	0.958	0.888	0.881	0.888			2×20	0.960	0.884	0.879	0.890
	Adaptive	2×5	0.795	0.784	0.783	0.754		Adaptive	2×5	0.514	0.514	0.501	0.473
		2×10	0.928	0.912	0.913	0.868			2×10	0.822	0.811	0.806	0.773
LINE (abrupt)		2×20	0.972	0.959	0.957	0.911	SINEH		2×20	0.889	0.874	0.877	0.833
(abrupt)	Non-adaptive	2×5	0.795	0.785	0.783	0.754	(abrupt)	Non-adaptive	2×5	0.514	0.514	0.503	0.474
		2×10	0.928	0.905	0.900	0.861			2×10	0.822	0.804	0.790	0.765
		2×20	0.972	0.953	0.887	0.883			2×20	0.889	0.862	0.822	0.808
LINE (gradual)	Adaptive	2×5	0.795	0.721	0.729	0.731		Adaptive	2×5	0.514	0.513	0.510	0.478
		2×10	0.928	0.870	0.880	0.855			2×10	0.822	0.756	0.770	0.756
		2×20	0.972	0.910	0.918	0.898	SINEH		2×20	0.889	0.812	0.820	0.818
	Non-adaptive	2×5	0.795	0.722	0.728	0.731	(graduar)	Non-adaptive	2×5	0.514	0.514	0.511	0.478
		2×10	0.928	0.871	0.879	0.855			2×10	0.822	0.763	0.774	0.757
		2×20	0.972	0.903	0.893	0.895			2×20	0.889	0.811	0.811	0.817

Table 3: Comparative results of sCA with and without adaptive mechanism with different grid sizes for synthetic CIRCLE, LINE, SINEV and SINEH datasets. The prequential accuracy is measured at points **c** (before the drift occurs), **d** (after drift occurs and adaptation mechanism has been triggered), and **e** (final). The last column amounts to the mean prequential accuracy averaged over the duration of the whole stream.

In light of the above claim, we can finally confirm that in case of abrupt drifts we should opt for small grid sizes G, which is better in terms of computational cost. However, for gradual drifts the grid should be made finer in order to emphasize the differences between adaptive and non-adaptive versions. For the sake of space, only the cases of CIRCLE and SINEH datasets are presented in Figures 2 (abrupt drift) and 3 (gradual drift), and 4 (abrupt drift) and 5 (gradual drift), respectively. The results for all the synthetic datasets can be found in Table 3.

5.3. RQ3: Is LUNAR competitive in comparison with other consolidated OLMs of the literature?

In Table 4 and Figure 6 we analyze the competitiveness of LUNAR in terms of classification performance for the considered real-world datasets. In the ELEC2 dataset, only SGDC (0.820) outperforms LUNAR (0.763), and HTC shows almost the same score (0.766). In case of the GMSC dataset, only HTC (0.910) and PAC (0.891) outperforms LUNAR (0.869). For the POKER-HAND dataset, only HTC (0.573) and KNNC (0.567) outperforms LUNAR (0.527).

Following the evaluation method used in (Bifet, Holmes, Pfahringer & Frank, 2010), the column *Global mean* preACC of Table 4 averages the results in all datasets, and shows how LUNAR (0.719) is the second best method, only surpassed by HTC (0.749), which is arguably one of the best stream learners in the field. After the analysis of the results, we can confirm that effectively LUNAR has turned into a very competitive stream learning method.

Method	Dataset	preACC (mean±std)	Global mean preACC
LUNAR	ELEC2 GMSC POKER-HAND	0.763 ± 0.0 0.869 ± 0.0 0.527 ± 0.0	$\boldsymbol{0.719\pm0.0}$
SGDC	ELEC2 GMSC POKER-HAND	0.820 ± 0.058 0.845 ± 0.085 0.446 ± 0.061	0.703 ± 0.068
HTC	ELEC2 GMSC POKER-HAND	0.766 ± 0.0 0.910 ± 0.0 0.573 ± 0.0	$\boldsymbol{0.749 \pm 0.0}$
PAC	ELEC2 GMSC POKER-HAND	$\begin{array}{c} 0.670 \pm 0.043 \\ 0.891 \pm 0.032 \\ 0.496 \pm 0.015 \end{array}$	0.685 ± 0.030
KNNC	ELEC2 GMSC POKER-HAND	$\begin{array}{c} 0.621 \pm 0.019 \\ 0.822 \pm 0.014 \\ 0.567 \pm 0.055 \end{array}$	0.670 ± 0.029

Table 4: Comparative results of LUNAR with other stream learners in real-world datasets. The column **preACC** (mean±std) denotes the mean prequential accuracy of every method in each dataset. The column **Global mean preACC** presents the mean prequential accuracy for all datasets of each method, and serves as a reference for a global comparison. Global results of the two best methods are marked in bold.

5.4. Final Observations, Remarks and Recommendations

Once the three research questions have been thoroughly answered, it is worth discussing some general recommendations related to LUNAR. When designing a sCA for non-stationary scenarios, we should consider that generally we will assign one grid dimension to each feature of the dataset, thus as many dimensions in the grid and as many cells per feature, as much computational cost and more processing time will be required. Specifically, provided that a problem with d dimensions is under target, and given a granularity (size) of the grid given by G, the worst-case complexity of predicting the class of a given test instance X_t is given by $\mathcal{O}(G^d)$, which is the time taken by a single processing thread to explore all cells of the d-dimensional grid of cells and discriminate the cell enclosing X_t . Due to the exponential complexity, we recommend the use of sCA (and CA in pattern recognition in general) in datasets with a low num-



Figure 6: Prequential accuracy preACC(t) for LUNAR and the OLMs under consideration in the selected real-world datasets. The first 50% of the dataset is for preparatory instances, while the rest is adopted as streaming *test-then-train* instances. A moving window of 500 instances has been applied in order to smooth out short-term fluctuations and to furnish a more friendly visualization.

ber of features. Nevertheless, the search process over the grid's cells can be easily parallelized, hence allowing for fast prediction and cell updating speeds.

We also pause at the *streamified* version of the KNNC model, whose similarity to LUNAR calls for a brief comparison among them. In essence, both learning models rely on the concept of neighborhood in the feature space, induced by either the selected measure of similarity among instances (KNNC) or the arrangement of cells in a grid (LUNAR). We have seen in Table 4 that our proposed method overcomes the results of KNNC in ELEC2 and GMSC datasets. Here it is worth mentioning that in order to have a fair comparison between both methods, the number of nearest neighbors to search for in the KNNC method has been properly optimized before the streaming process runs. Furthermore, the maximum size of the window storing the last viewed instances of KNNC has been set equal to the W parameter in LUNAR. The capability of the sCA grid's cells to reflect and maintain the prevailing knowledge in the stream renders a superior performance than that offered by the KNNC method when computing the similarity over the W-sized window of past instances.

6. Conclusions and Future Work

Cellular automata have been successfully applied to different real-world applications since their inception several decades ago. Among them, pattern recognition has been proven to be a task in which the self-organization and modeling capabilities of cellular automata can yield significant performance gains. This work has built upon this background to explore whether such benefits can also be extrapolated to real-time machine learning under non-stationary conditions, which is arguably among the hottest topics in Data Science nowadays. Under the premises of real-time settings, aspects such as complexity and adaptability to changing tasks are also important modeling design drivers that add to the performance of the model itself.

The algorithm introduced in this work provides a new perspective in the stream learning scene. We have proposed a cellular automaton able to learn incrementally and capable of adapting to evolving environments (LUNAR), showing a competitive classification performance when is compared with other reputed state-of-the-art algorithms. LUNAR contributes to the discussion on ways to use cellular automata for paradigms in which the computation effort relies on a network of simple, interconnected devices with very low processing capabilities and constrained battery capacity (e.g. Utility Fog, Smart Dust, MEMS or Swarms). Under these circumstances, learning algorithms should be embarked in miniaturized devices running on low-power hardware with limited storage.

LUNAR has shown a well performance in practice over the datasets considered in this study, with empirical evidences of its adaptability when mining non-stationary data streams. Future work will be devoted towards experimenting with other local rules or neighborhoods to determine their effects on these identified properties of cellular automata. Some preliminary experiments carried out off-line suggest that a sCA could also *detect* drifts, which, along with their simplicity, paves the way towards adopting

them in active strategies for stream learning in non-stationary setups. We have also conceived the possibility of configuring ensembles of sCA, wherein diversity among the constituent automata can be induced by very diverse means (e.g. online bagging, boosting or probabilistic class switching). We strongly believe that moving in this algorithmic direction may open up the chance of designing more powerful cellular automata with complementary, potentially better capabilities to deal with stream learning scenarios.

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